



## **A tale of genome, annotations, metabolism and phylogenomics: the pea aphid genome resources**

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# A tale of genome, annotations, metabolism and phylogenomics: the pea aphid genome resources

Fabrice Legeai<sup>1,2</sup>, Stefano Colella<sup>3,4</sup>, Jaime Huerta-Cepas<sup>5</sup>, J-P Gauthier<sup>1</sup>, Augusto Vellozo<sup>4,6</sup>, Patrice Baa-Puyoulet<sup>3</sup>, Marie-France Sagot<sup>4,6</sup>,

Toni Gabaldon<sup>5</sup>, Olivier Collin<sup>2</sup>, Hubert Charles<sup>3,4</sup>, Denis Tagu<sup>1</sup>

<sup>1</sup> INRA, BiO3P, Rennes, 35653, France, <sup>2</sup> INRIA-IRISA, Centre Rennes-Bretagne-Atlantique, Rennes, 35000, France, <sup>3</sup> INRA-INSA, BF2I, Villeurbanne, 69100, France,

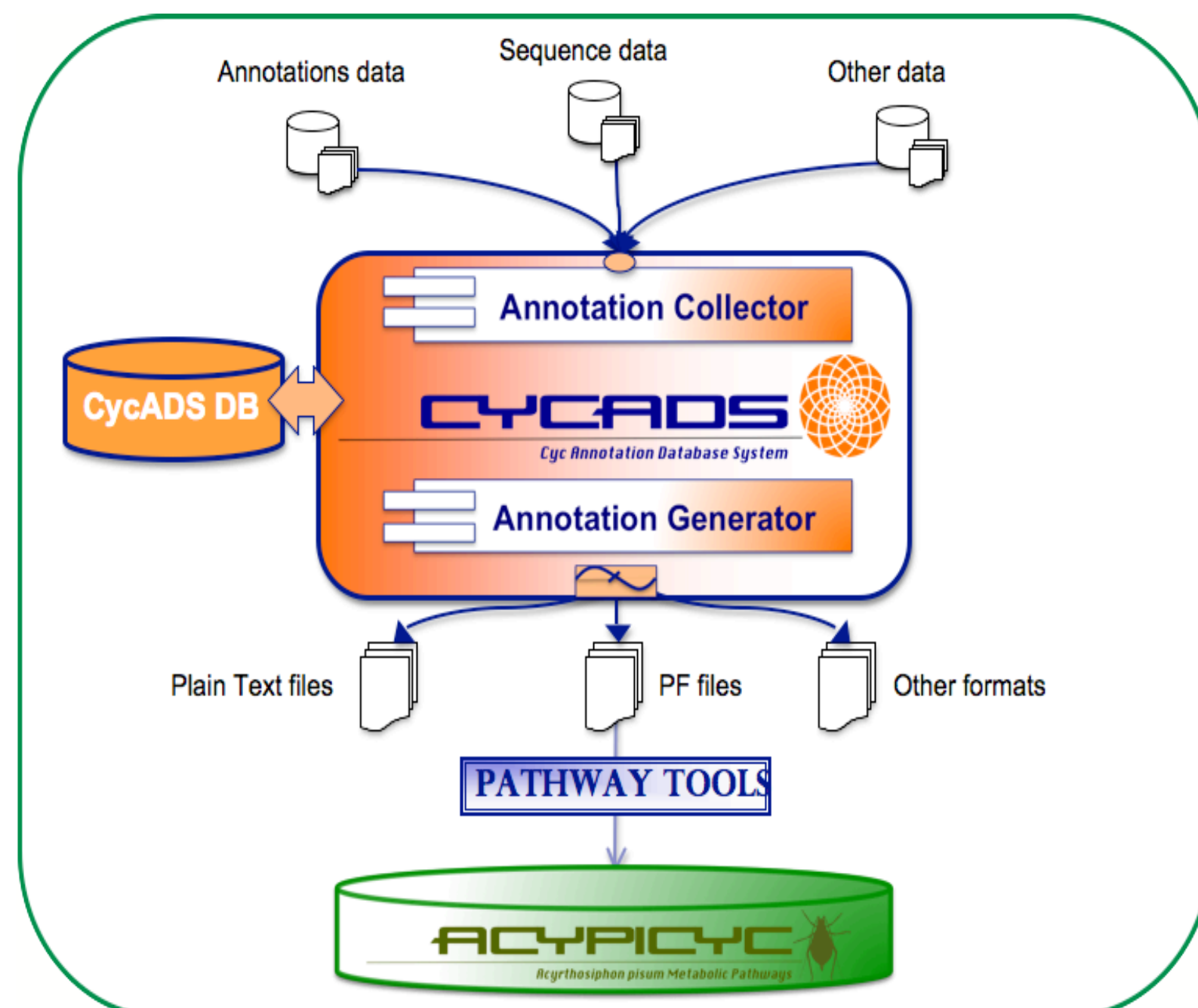
<sup>4</sup> INRIA, Bamboo, Lyon, 69000, France, <sup>5</sup> CRG, Centre for Genomic Regulation, Barcelona, 08003, Spain, <sup>6</sup> UCBL1, LBBE, Villeurbanne, 69100, France



## CycADS: the Cyc Annotation Database System

The CycADS pipeline has proven to be useful in the generation of the AcyciCyc database and we plan to use the same metabolism genes annotation strategy for other arthropods sequenced genomes.

**WORKFLOW : from CycADS to AcyciCyc, and beyond...**



Data from GenBank and different gene annotation tools (such as KAAS, PRIAM, Blast2GO, PhylomeDB, etc, ...) are collected in an *ad hoc* SQL database, the core component of CycADS

A set of Java programs allows the data upload from the different annotation sources.

Each annotation receives an evidence score and a specific filter is applied to extract the best annotation that is then included in the "Pathologic" file (PF) used by Pathway Tools to generate a filtered and enriched Cyc database.

The CycADS pipeline allows automated updates of a given BioCyc database as soon as new gene/protein annotation data are available

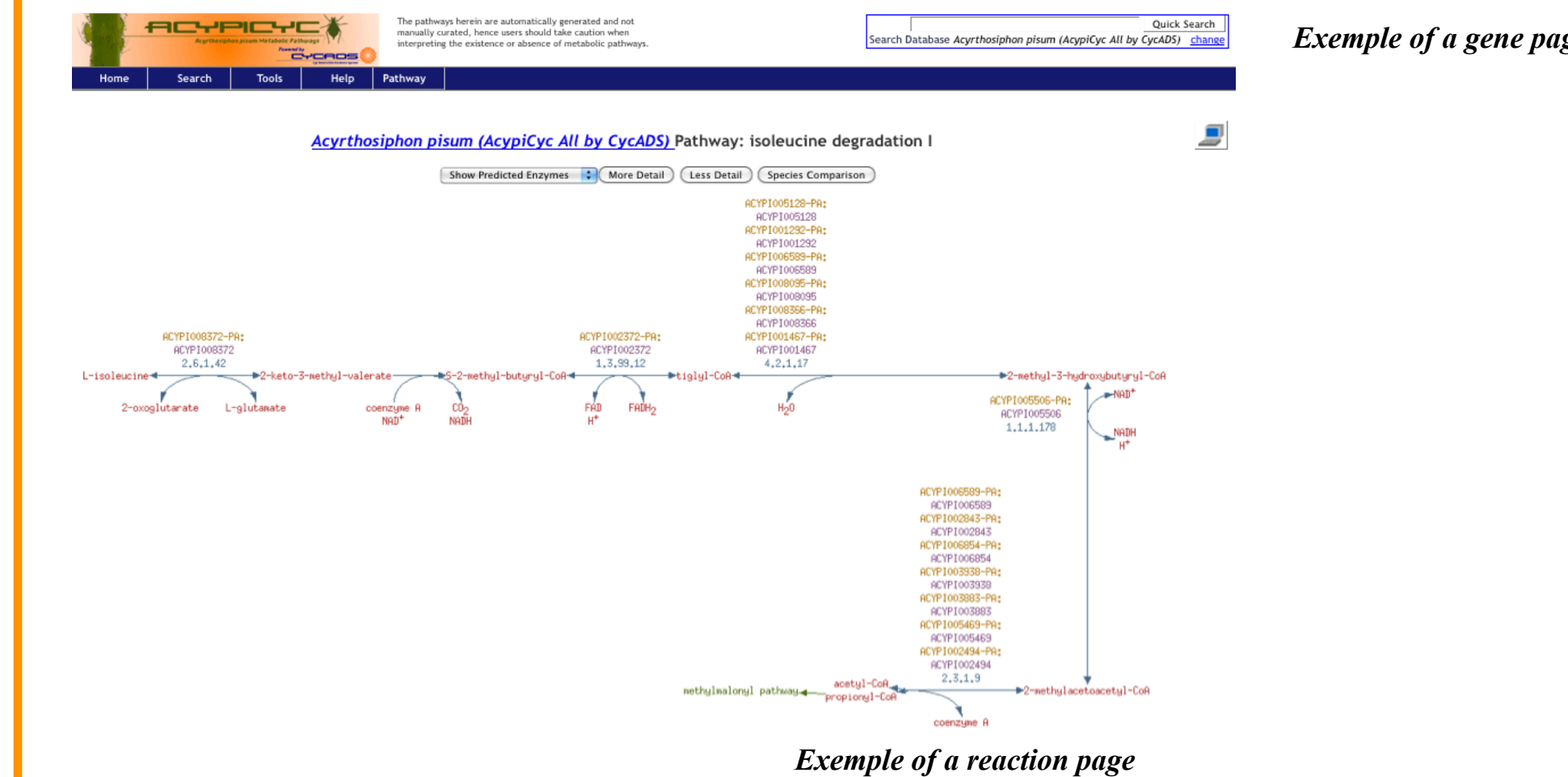
## AcyciCyc: a CycADS powered database

AcyciCyc, as all BioCyc databases, offers a framework for the analysis of the integrated metabolic network and different query tools allow the user to visualize different metabolic reactions and pathways. Thanks to CycADS several supplementary specific links can be added to complement the classic existing ones. This feature is most valuable for newly sequenced genomes that are kept in community based repository (such as AphidBase for the pea aphid).

Not only enzymes, but all genes are present in AcyciCyc. All gene pages include an annotation summary with an associated score and a set of hyperlinks to different information resources including genomics (AphidBase and GenBank), phylogenomics (PhylomeDB) and metabolism (KEGG orthology, BRENDA, ENZYME) database



Example of a gene page

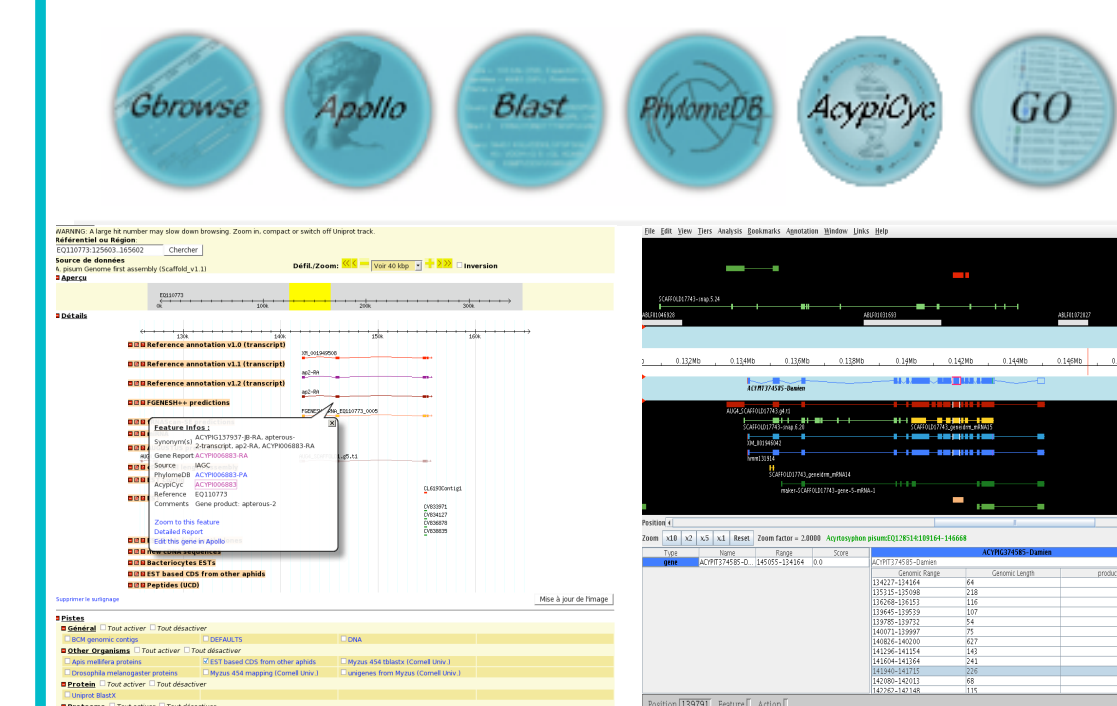


Example of a reaction page

**URLs**  
AcyciCyc : <http://acycicyc.cycadsys.org/>  
CycADS : [http://www4.inra.fr/cycads\\_eng](http://www4.inra.fr/cycads_eng)

## AphidBase : a GMOD Chado database

AphidBase is a comprehensive information system set up to safely centralize, manage, mine, disseminate and promulgate data generated by the International Aphid Genomics Consortium.



This system was built using open-source software tools from GMOD including several Chado instances, a genome browser (Gbrowse), Apollo for the manual curation, and various other tools such as a blast search and a full text search facilities.

It allowed an international broad community dispersed at many sites to produce a robust and comprehensive annotation of the pea aphid genome by curating gene models and gathering manual and functional annotations, which is an essential step to attain a basal data quality.

Version	gene predictions	RefSeq Models	Glean Models	Total
Acypa v1.0	10466	24395	34821	
Acypa v1.1	Appraised mRNAs	1,255 (12.0%)	168 (0.7%)	1,423 (4.1%)
	Corrected mRNAs	231 (18.4%)	125 (74.4%)	356 (25.0%)
	Validated mRNAs	1024	43	1067
	New genes	141	141	282
	unchecked RNAs	9211	24187	33398
Acypa v1.2	Appraised mRNAs	1,297 (12.4%)	329 (1.3%)	1,626 (4.7%)
	Corrected mRNAs	284 (21.9%)	286 (87.0%)	570 (16.6%)
	Validated mRNAs	1,013	43	1,056
	New genes	300	300	600
	unchecked RNAs	9,164	24,027	33,191
Others	miRNA	189	384	573
	rRNA	384	263,048	263,432
	A. pisum EST/unigene mapping	30,840	30,840	61,680
	Peptide alignments	236,670	236,670	473,340
	Transposable Elements locations	498,474	498,474	996,948

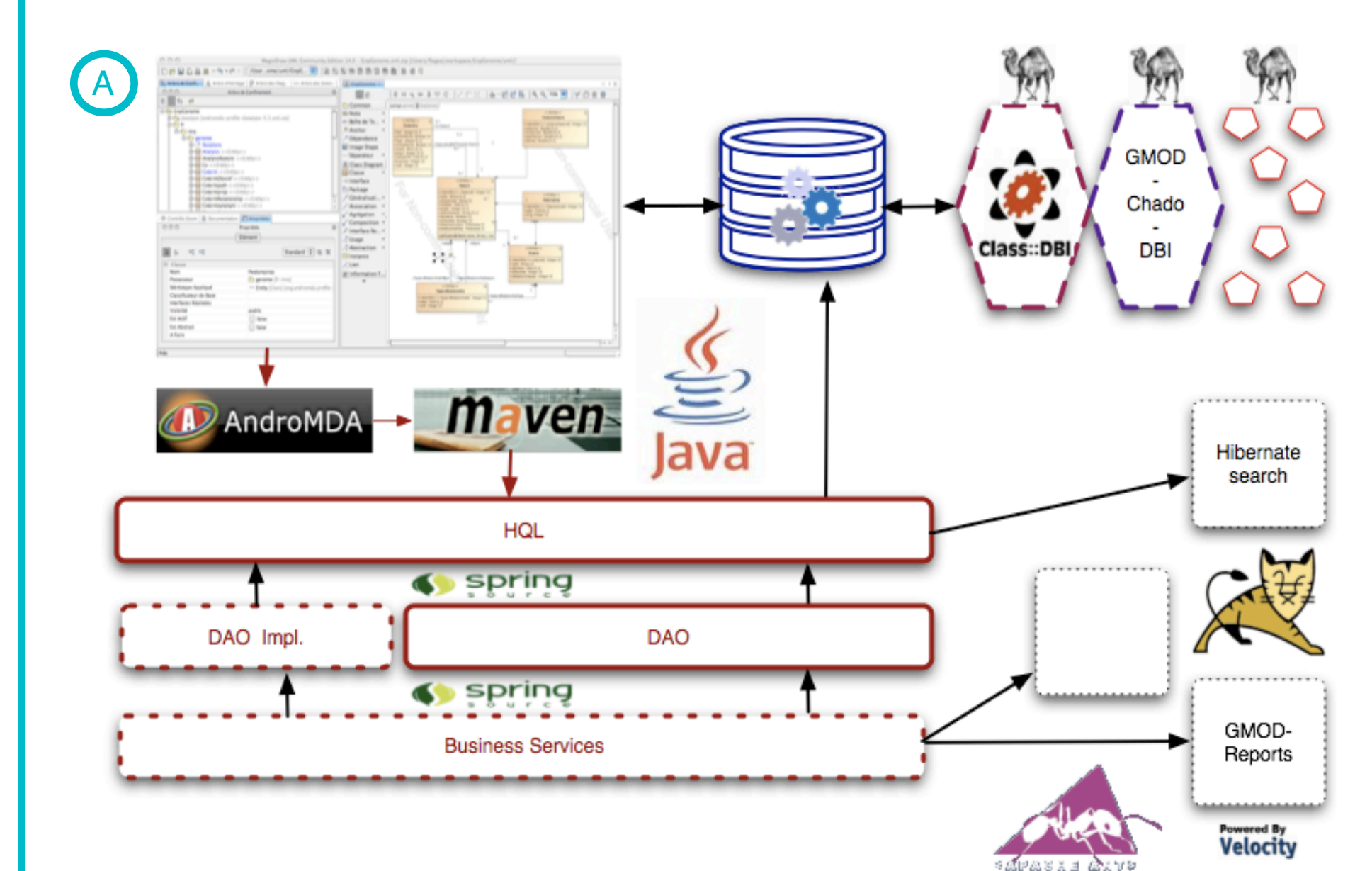
table of the main features stored in AphidBase

This system will be extended to support complete genomes sequencing or resequencing projects, and other projects based on deep sequencing strategies (expression profiling by RNA-Seq, variability studies, ChIP-Seq, ...) for various insect pest species. And, to ensure that all these resources were fully exploited by the community, it would be accompanied by an **AphidAtlas** aiming to link morphological characterizations of the aphids to transcriptomics data.

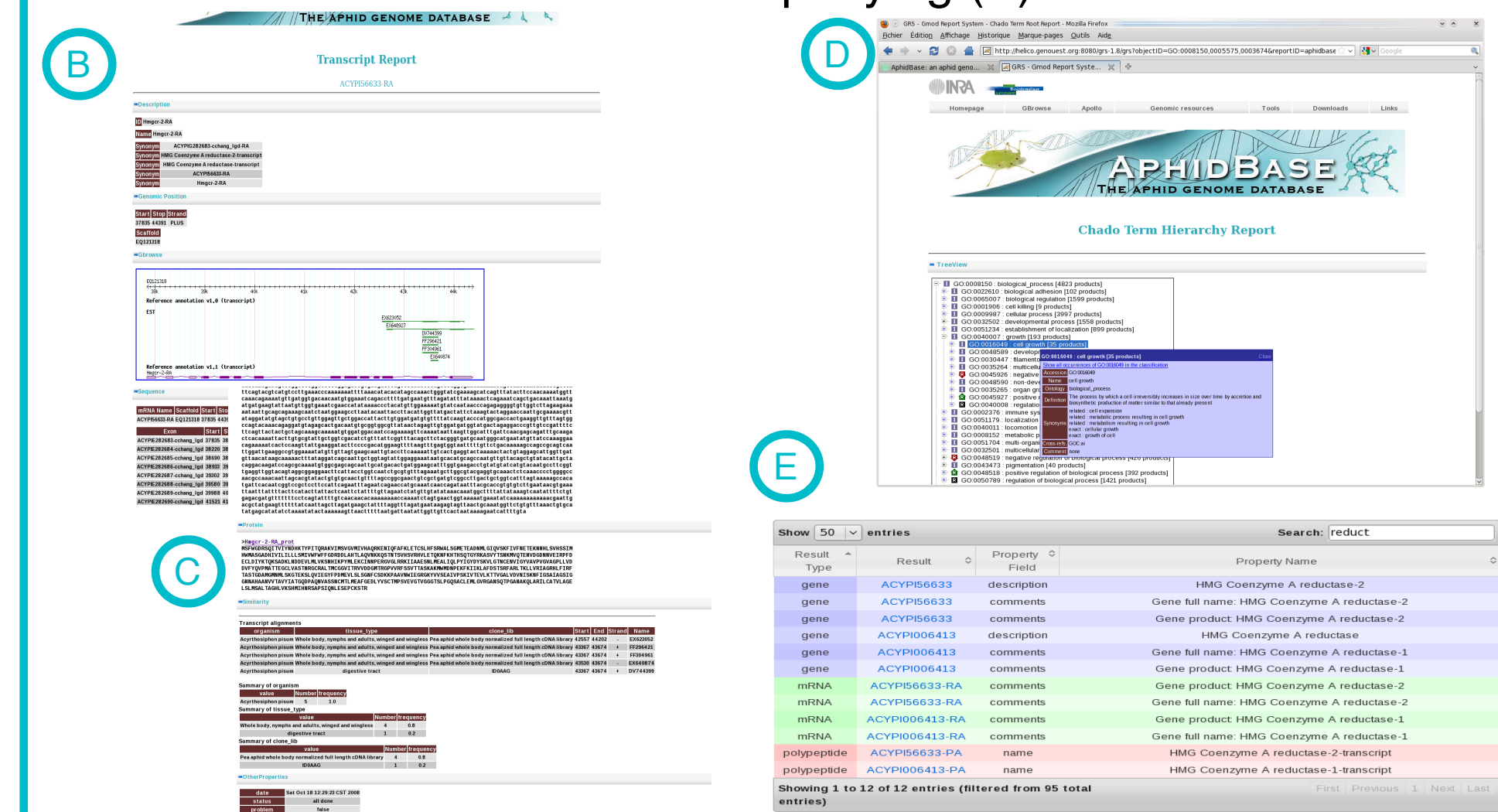
URL: <http://www.aphidbase.org>

## Chado Module : a Java middleware for Chado

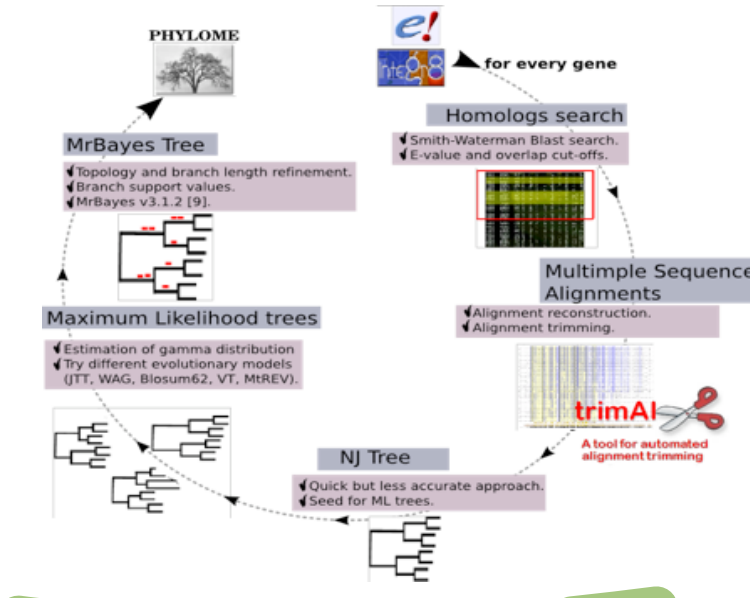
For connecting applications or scripts handling or presenting data stored into our GMOD-Chado databases, we developed two middleware, in Perl and in Java.



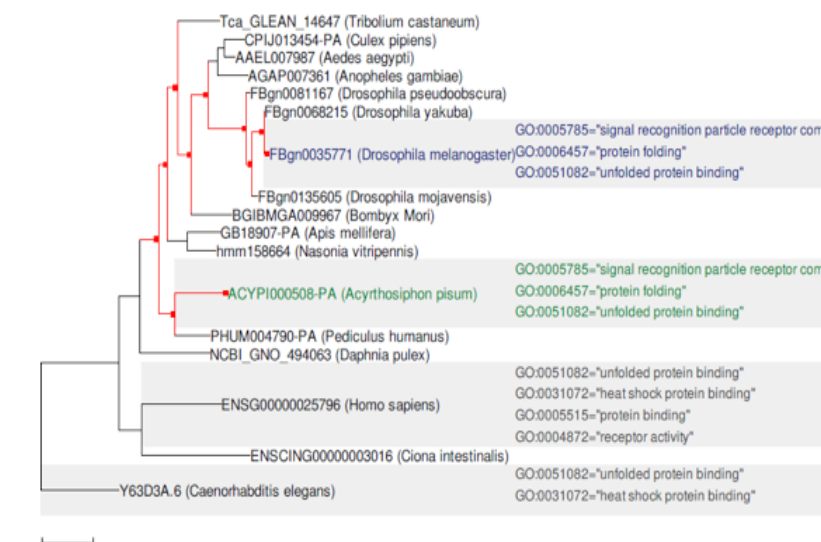
We used JAVA I2EE and a model-driven-architecture system (AndroMDA) for automatically generating the Hibernate classes from a Chado UML (A). Using those layers, we implemented 2 applications deployed on a Tomcat server : GMD reports, a configurable reports builder for browsing gene, transcript or proteins (B,C) and navigating into the ontologies (D), and Chado search for accelerated database querying (E).



A genome based phylogenetic analysis was performed for the pea aphid using the Phylome pipeline and the results are stored in the PhylomeDB database. Links to the database are provided both from AcyciCyc and AphidBase



The phylome analysis has also allowed the transfer of GO annotation from Drosophila melanogaster to the pea aphid.



ETE: Environment for Tree Exploration  
[ete.cgenomics.org](http://ete.cgenomics.org)  
Parallelization of PhyML in graphical cards (with O. Gascuel)

Huerta-Cepas et al.  
Nucleic Acids Res. (2008)  
[www.phylomedb.org](http://www.phylomedb.org)